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QIIME2

Anticipated workshop duration when delivered to a group of participants is 4 hours.

For queries relating to this workshop, contact Melbourne Bioinformatics (bioinformatics-training@unimelb.edu.au).

Overview

Topic

- Genomics
- Transcriptomics
- Proteomics

- Metabolomics
- Statistics and visualisation
- Structural Modelling
- Basic skills

Skill level

- Beginner
- Intermediate
- Advanced

This workshop is designed for participants with command-line knowledge. You will need to be able to `ssh` into a remote machine, navigate the directory structure and `scp` files from a remote computer to your local computer.

Description

What is the influence of genotype (intrinsic) and environment (extrinsic) on anemone-associated bacterial communities?

Data: Illumina MiSeq v3 paired-end $(2 \times 300 \text{ bp})$ reads (FASTQ).

Tools: QIIME 2

Pipeline:

Section 1: Importing, cleaning and quality control of the data

Section 2: Taxonomic Analysis

Section 3: Building a phylogenetic tree

Section 4: Basic visualisations and statistics

Section 5: Exporting data for further analysis in R

Learning Objectives

At the end of this introductory workshop, you will:

- Take raw data from a sequencing facility and end with publication quality graphics and statistics
- Answer the question What is the influence of genotype (intrinsic) and environment (extrinsic) on anemone-associated bacterial communities?

Tutorial layout

• There is a Table of contents on the right-hand side which can be used to easily navigate through the tutorial by clicking the relevant section.

These grey coloured boxes are code blocks. The rectangular boxes in the top

right hand corner of this code block/grey box can be used to copy the code to the clipboard.

??? example "Coloured boxes like these with > on the far right hand side, can be clicked to reveal the contents." REVEALED!

!!! attention "Attention: Pay attention to the information in these boxes." Important information, hints and tips.

Requirements and preparation

!!! attention "Important" **Attendees are required to use their own laptop computers.**

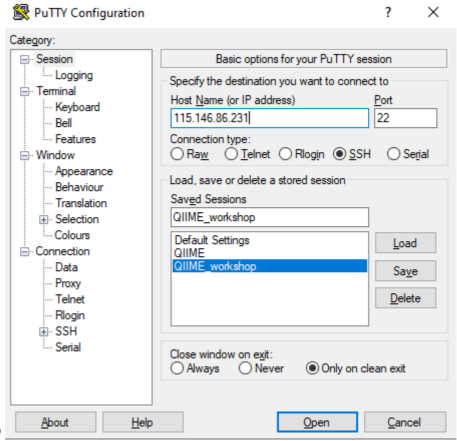
At least one week before the workshop, if required, participants should install the software below. This should provide sufficient time for participants to liaise with their own IT support should they encounter any IT problems.

Required Software

Mac Users: No additional software needs to be installed for this workshop.

Windows Users:

1. A terminal emulator such as PuTTY (free and open-source) will need to be downloaded.



??? example "Putty Example"

2. Software for file transfers between a local computer and remote server such as WinSCP or FileZilla.

Mode of Delivery

This workshop will be run on a Nectar Instance. An "Instance" is Nectar terminology for a virtual machine running on the Nectar Cloud OpenStack infrastructure. An "Instance" runs on a "compute node"; i.e. a physical computer populated with processor chips, memory chips and so on.

You will be given an individual username, IP address and password to log on to using the SSH client tool on your computer (Terminal on Mac or PuTTY on Windows).

ssh username@nectar_ip-address

Should you wish to do this tutorial at a later stage independently, it is possible to apply for your own instance directly through a [Nectar allocation]

(https://support.ehelp.edu.au/support/solutions/articles/6000068044-managing-an-allocation). There are also many helpful [Nectar Research Cloud tutorials](https://tutorials.rc.nectar.org.au/).

Byobu-screen

Some of the commands in this tutorial take a while to run. Should your connection drop and the SSH session on Nectar terminates, any commands that are running will terminate too. To mitigate this, once logged on to the Nectar Instance, we'll run byobu—screen (an enhancement for the screen terminal

multiplexer) which allows us to resume a session. In other words, processes running in byobu—screen will continue to run when their window is not visible, even if you get disconnected.

On Nectar, to start a byobu-screen session called workshop, type

byobu-screen -S workshop

??? example "Byobu Example"

(qiime2-2021.4) epsilon@test-i5:~\$



You can then proceed to run the commands in the workshop as normal.

Should your SSH session on Nectar terminate, once you log back in to your Nectar instance, list running sessions/screens:

byobu-screen -ls

If it says (Detached) next to the workshop session in the list, reattach to workshop by:

byobu-screen -r workshop

If it says (Attached) next to the workshop session in the list, you can access workshop which is already attached by:

```
byobu-screen -r -d workshop
```

Some other useful 'byobu-screen' commands:

- To detach from workshop, type ctrl-a ctrl-d while inside the workshop session. (You will need to configure Byobu's ctrl-a behaviour if it hasn't already been configured (text will appear on the screen telling you this). Follow the information on the screen and select 1 for Screen mode).
- To terminate workshop, type ctrl-d while inside the workshop session.

Required Data

- No additional data needs to be downloaded for this workshop it is all located on the Nectar
 Instance. FASTQs are located in the directory raw_data and a metadata (metadata.tsv) file has
 also been provided.
- If you wish to analyse the data independently at a later stage, it can be downloaded from here. This zipped folder contains both the FASTQs and associated metadata file.

Symbolic links to workshop data

Data for this workshop is stored in a central location (/mnt/shared_data/) on the Nectar file system that we will be using. We will use symbolic links (ln -s) to point to it. Symbolic links (or symlinks) are just "virtual" files or folders (they only take up a very little space) that point to a physical file or folder located elsewhere in the file system. Sequencing data can be large, and rather than unnecessarily having multiple copies of the data which can quickly take up a lot of space, we will simply point to the files needed in the shared_data folder.

```
cd
ln -s /mnt/shared_data/raw_data raw_data
ln -s /mnt/shared_data/metadata.tsv metadata.tsv
ln -s /mnt/shared_data/silva_138_16s_v5v6_classifier_2021-4.qza
silva_138_16s_v5v6_classifier_2021-4.qza
```

Slides and workshop instructions

Click here for slides presented during this workshop.

Click here for a printer friendly PDF version of this workshop.

Author Information

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Created/Reviewed: August 2021

Background

What is the influence of genotype (intrinsic) and environment (extrinsic) on anemone-associated bacterial communities?

The Players

- Exaiptasia diaphana a shallow-water, marine anemone that is often used in research as a model organism for corals. In this experiment, two genotypes (AIMS1 and AIMS4) of *E. diaphana* were grown in each of two different environments:
 - 1. sterile seawater OR
 - 2. unfiltered control seawater
- The anemone-associated bacterial communities or *microbiome* these bacteria live on, or within *E. diaphana*, and likely consist of a combination of commensals, transients, and long-term stable members, and combined with their host, form a mutually beneficial, stable symbiosis.

The Study

The anemone microbiome contributes to the overall health of this complex system and can evolve in tandem with the anemone host. In this data set we are looking at the impact of intrinsic and extrinsic factors on anemone microbiome composition. After three weeks in either sterile or control seawater (environment), anemones were homogenized and DNA was extracted. There are 23 samples in this data set - 5 from each anemone treatment combination (2 genotypes x 2 environments) and 3 DNA extraction blanks as controls. This data is a subset from a larger experiment.

Dungan AM, van Oppen MJH, and Blackall LL (2021) Short-Term Exposure to Sterile Seawater Reduces Bacterial Community Diversity in the Sea Anemone, *Exaiptasia diaphana*. *Front. Mar. Sci.* 7:599314. doi:10.3389/fmars.2020.599314 [Full Text].

QIIME 2 Analysis platform

Quantitative Insights Into Microbial Ecology 2 (QIIME 2^{TM}) is a next-generation microbiome bioinformatics platform that is extensible, free, open source, and community developed. It allows researchers to:

- Automatically track analyses with decentralised data provenance
- Interactively explore data with beautiful visualisations
- · Easily share results without QIIME 2 installed
- Plugin-based system researchers can add in tools as they wish

!!! attention The version used in this workshop is qiime2-2021.4. Other versions of QIIME2 may result in minor differences in results.

Viewing QIIME2 visualisations

As this workshop is being run on a remote Nectar Instance, you will need to download the visual files (*.qzv) to your local computer and view them in QIIME 2 View (q2view).

!!! attention We will be doing this step multiple times throughout this workshop to view visualisation files as they are generated.

Mac Users

The syntax to do this depends on whether you are running the copying command on your local computer, or on the remote computer (Nectar cloud).

1. When running the command from your local computer, the syntax for copying a file from Nectar is:

```
scp username@nectar_IP_address:FILENAME /PATH/TO/TARGET/FOLDER/
```

2. Running the command on the remote computer, the syntax for copying a file *to* your local computer is:

```
scp FILENAME username@your_IP_address:/PATH/TO/TARGET/FOLDER/
```

Less experienced Unix users may want to use FileZilla.

Windows Users

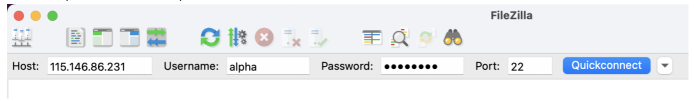
Using WinSCP or FileZilla:

Host: The IP address of the Nectar instance

Username: alpha | beta | gamma | delta | epsilon | delta

Port: 22

??? example "Filezilla Example"



Alternatively, ***if you have QIIME2 installed and are running it on your own computer***, you can use `qiime tools view` to view the results from the command line (e.g. `qiime tools view filename.qzv`). `qiime tools view` opens a browser window with your visualization loaded in it. When you are done, you can close the browser window and press `ctrl-c` on the keyboard to terminate the command.

Section 1: Importing, cleaning and quality control of the data

Import data

These samples were sequenced on a single Illumina MiSeq run using v3 (2 × 300 bp) reagents at the Walter and Eliza Hall Institute (WEHI), Melbourne, Australia. Data from WEHI came as paired-end, demultiplexed, unzipped *.fastq files with adapters still attached. Following the QIIME2 importing tutorial, this is the Casava One Eight format. The files have been renamed to satisfy the Casava format as SampleID_FWDXX-REVXX_L001_R[1 or 2]_001.fastq e.g. CTRLA_Fwd04-Rev25_L001_R1_001.fastq.gz. The files were then zipped (.gzip).

Here, the data files (two per sample i.e. forward and reverse reads R1 and R2 respectively) will be imported and exported as a single QIIME 2 artefact file. These samples are already demultiplexed (i.e. sequences from each sample have been written to separate files), so a metadata file is not initially required.

!!! note To check the input syntax for any QIIME2 command, enter the command, followed by --help e.g.
qiime tools import --help

!!! attention If you haven't already done so, make sure you are running the workshop in byobu-screen and have created the symbolic links to the workshop data.

Start by making a new directory analysis to store all the output files from this tutorial. In addition, we will create a subdirectory called seqs to store the exported sequences.

```
cd
mkdir —p analysis/seqs
```

Run the command to import the raw data located in the directory raw_data and export it to a single QIIME 2 artefact file, combined.qza.

```
qiime tools import \
   --type 'SampleData[PairedEndSequencesWithQuality]' \
   --input-path raw_data \
   --input-format CasavaOneEightSingleLanePerSampleDirFmt \
   --output-path analysis/seqs/combined.qza
```

Remove primers

!!! important Remember to ask you sequencing facility if the raw data you get has the primers attached - they may have already been removed.

These sequences still have the primers attached - they need to be removed (using cutadapt) before denoising.

```
qiime cutadapt trim-paired \
  --i-demultiplexed-sequences analysis/seqs/combined.qza \
```

```
--p-front-f AGGATTAGATACCCTGGTA \
--p-front-r CRRCACGAGCTGACGAC \
--p-error-rate 0.20 \
--output-dir analysis/seqs_trimmed \
--verbose
```

!!! attention The primers specified (784f and 1492r for the bacterial 16S rRNA gene) correspond to *this* specific experiment - they will likely not work for your own data analyses.

!!! attention The error rate parameter, #!python --p-error-rate, will likely need to be adjusted for your own sample data to get 100% (or close to it) of reads trimmed.

Create and interpret sequence quality data

Create a viewable summary file so the data quality can be checked. Viewing the quality plots generated here helps determine trim settings.

!!! info "Things to look for:" 1. Where does the median quality drop below 30?

2. Do any of the samples have only a few sequences e.g. <1000? If so, you may want to omit them from the analysis later on in R.

Create a subdirectory in analysis called visualisations to store all files that we will visualise in one place.

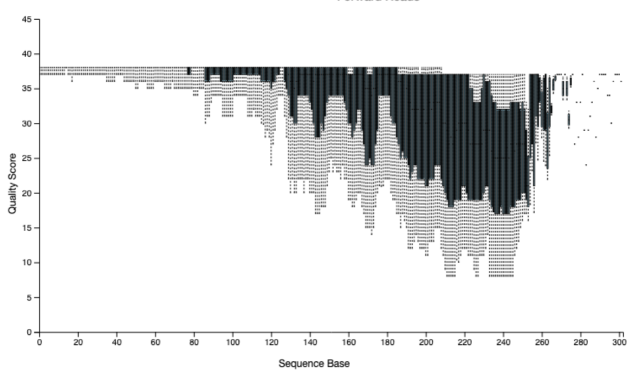
```
mkdir analysis/visualisations
```

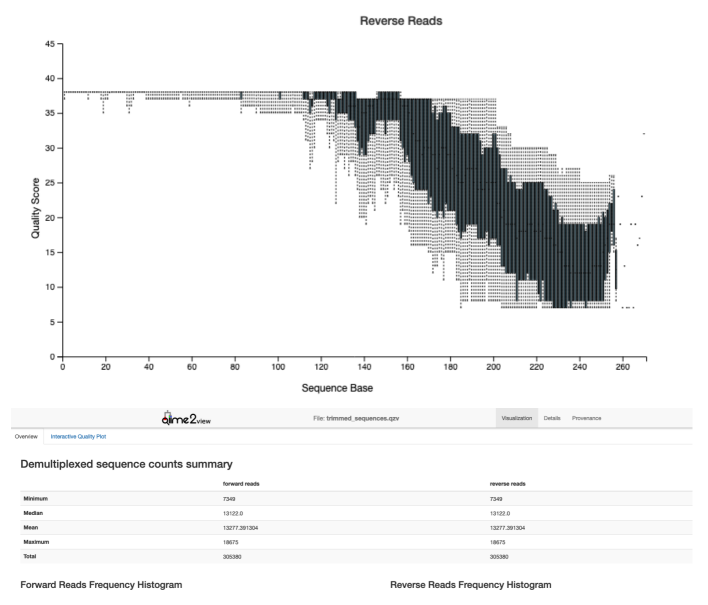
```
qiime demux summarize \
   --i-data analysis/seqs_trimmed/trimmed_sequences.qza \
   --o-visualization analysis/visualisations/trimmed_sequences.qzv
```

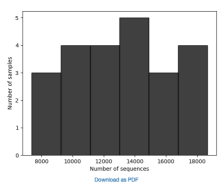
Copy analysis/visualisations/trimmed_sequences.qzv to your local computer and view in QIIME 2 View (q2view).

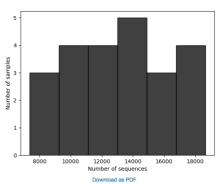
??? example "Visualisations: Read quality and demux output"

Forward Reads









Denoising the data

Trimmed sequences are now quality assessed using the dada2 plugin within QIIME2. dada2 denoises data by modelling and correcting Illumina-sequenced amplicon errors, and infers exact amplicon sequence variants (*ASVs*), resolving differences of as little as 1 nucleotide. Its workflow consists of filtering, dereplication, reference-free chimera detection, and paired-end reads merging, resulting in a feature or *ASV* table.

!!! note This step may long time to run (i.e. hours), depending on files sizes and computational power.

```
Remember to adjust `p-trunc-len-f` and `p-trunc-len-r` values according to your own data.
```

!!! question "Question: Based on your assessment of the quality plots from the trimmed_sequences.qzv file generated in the previous step, what values would you select for p-trunc-len-f and p-trunc-len-r in the command below?"

```
??? answer
  `p-trunc-len-f 211` and `p-trunc-len-r 172`
```

The specified output directory must not pre-exist.

```
qiime dada2 denoise-paired \
    --i-demultiplexed-seqs analysis/seqs_trimmed/trimmed_sequences.qza \
    --p-trunc-len-f xx \
    --p-trunc-len-r xx \
    --p-n-threads 0 \
    --output-dir analysis/dada2out \
    --verbose
```

Generate summary files

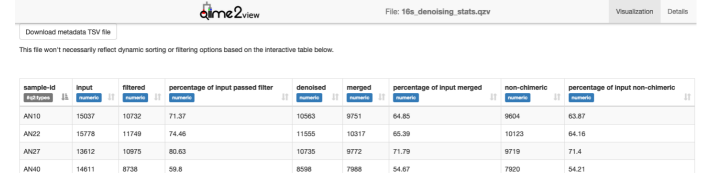
A metadata file is required which provides the key to gaining biological insight from your data. The file metadata.tsv is provided in the home directory of your Nectar instance. This spreadsheet has already been verified using the plugin for Google Sheets, keemei.

!!! info "**Things to look for:**" 1. How many features (*ASVs*) were generated? Are the communities high or low diversity? 2. Do BLAST searches of the representative sequences make sense? Are the features what you would expect e.g. marine or terrestrial? 3. Have a large number (e.g. >50%) of sequences been lost during denoising/filtering? If so, the settings might be too stringent.

```python qiime metadata tabulate \ --m-input-file analysis/dada2out/denoising\_stats.qza \ --o-visualization analysis/visualisations/16s\_denoising\_stats.qzv \ --verbose ```

Copy analysis/visualisations/16s\_denoising\_stats.qzv to your local computer and view in QIIME 2 View (q2view).

## ??? example "Visualisation: Denoising Stats"



9790

7332

9020

53.83

52.06

76.57

```python qiime feature-table summarize \ --i-table analysis/dada2out/table.qza \ --m-sample-metadata-file metadata.tsv \ --o-visualization analysis/visualisations/16s\_table.qzv \ --verbose ```

7534

9104

77.28

10870

8298

9497

9063

Copy analysis/visualisations/16s_table.qzv to your local computer and view in QIIME 2 View (q2view).

??? example "Visualisations: Feature/ASV summary"

59.96

81.46

85.54

dime2 _{view}			File: 16s_table.qzv
Overview	Interactive Sample Detail	Feature Detail	

Table summary

AN45

AN48

AN58

AN66

18187

14083

11780

11046

8444

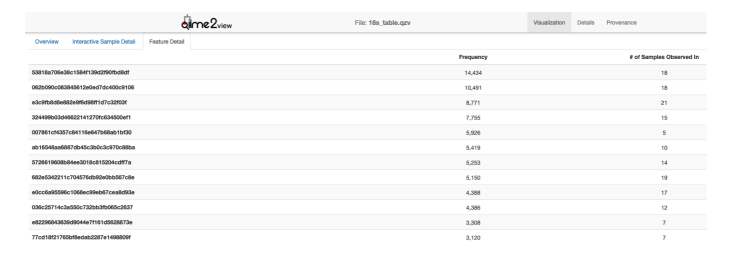
9596

Metric	Sample
Number of samples	23
Number of features	554
Total frequency	213,035

Frequency per sample

	Frequency
Minimum frequency	5,583.0
1st quartile	7,626.0
Median frequency	9,020.0
3rd quartile	10,137.5
Maximum frequency	14,021.0
Mean frequency	9,262.391304347826

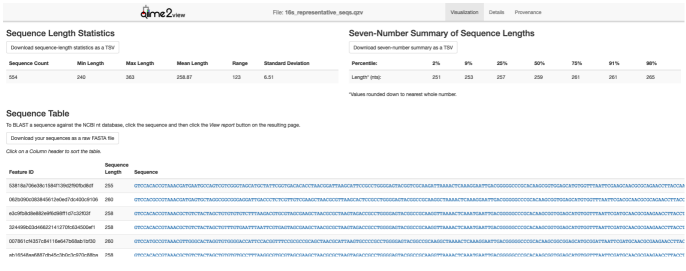
Frequency per sample detail (csv | html)



```python qiime feature-table tabulate-seqs \ --i-data analysis/dada2out/representative\_sequences.qza \ -- o-visualization analysis/visualisations/16s\_representative\_seqs.qzv \ --verbose ```

Copy analysis/visualisations/16s\_representative\_seqs.qzv to your local computer and view in QIIME 2 View (q2view).

??? example "Visualisation: Representative Sequences"



# Section 2: Taxonomic Analysis

## Assign taxonomy

Here we will classify each identical read or *Amplicon Sequence Variant (ASV)* to the highest resolution based on a database. Common databases for bacteria datasets are Greengenes, SILVA, Ribosomal Database Project, or Genome Taxonomy Database. See Porter and Hajibabaei, 2020 for a review of different classifiers for metabarcoding research. The classifier chosen is dependent upon:

- 1. Previously published data in a field
- 2. The target region of interest
- 3. The number of reference sequences for your organism in the database and how recently that database was updated.

A classifier has already been trained for you for the V5V6 region of the bacterial 16S rRNA gene using the SILVA database. The next step will take a while to run. *The output directory cannot previously exist*.

n\_jobs = 1 This runs the script using all available cores

!!! note The classifier used here is only appropriate for the specific 16S rRNA region that *this* data represents. It has also been trained specifically for the QIIME2 version that we are using in this workshop. You will need to train your own - no worries, QIIME2 has a tutorial for that.

```
qiime feature-classifier classify-sklearn \
 --i-classifier silva_138_16s_v5v6_classifier_2021-4.qza \
 --i-reads analysis/dada2out/representative_sequences.qza \
 --p-n-jobs 1 \
 --output-dir analysis/taxonomy \
 --verbose
```

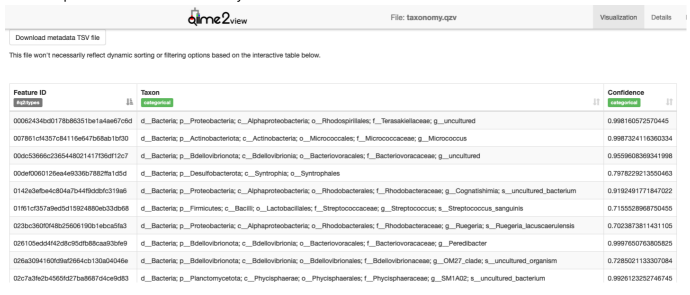
!!! warning "Warning" This step often runs out of memory on full datasets. Some options are to change the number of cores you are using (adjust --p-n-jobs) or add --p-reads-per-batch 10000 and try again. The QIIME 2 forum has many threads regarding this issue so always check there was well.

Generate a viewable summary file of the taxonomic assignments.

```
qiime metadata tabulate \
 --m-input-file analysis/taxonomy/classification.qza \
 --o-visualization analysis/visualisations/taxonomy.qzv \
 --verbose
```

Copy analysis/visualisations/taxonomy.qzv to your local computer and view in QIIME 2 View (q2view).

??? example "Visualisation: Taxonomy"



## **Filtering**

Filter out reads classified as mitochondria and chloroplast. Unassigned ASVs are retained. Generate a viewable summary file of the new table to see the effect of filtering.

According to QIIME developer Nicholas Bokulich, low abundance filtering (i.e. removing ASVs containing very few sequences) is not necessary under the ASV model.

```
qiime taxa filter-table \
 --i-table analysis/dada2out/table.qza \
 --i-taxonomy analysis/taxonomy/classification.qza \
 --p-exclude Mitochondria,Chloroplast \
 --o-filtered-table analysis/taxonomy/16s_table_filtered.qza \
 --verbose
```

```
qiime feature-table summarize \
 --i-table analysis/taxonomy/16s_table_filtered.qza \
 --m-sample-metadata-file metadata.tsv \
 --o-visualization analysis/visualisations/16s_table_filtered.qzv \
 --verbose
```

Copy analysis/visualisations/16s\_table\_filtered.qzv to your local computer and view in QIIME 2 View (q2view).

??? example "Visualisation: 16s\_table\_filtered"

dime			dime2view	File: 16s_table_filtered.qzv
	Overview	Interactive Sample Detail	Feature Detail	

# Table summary

Metric	Sample
Number of samples	23
Number of features	548
Total frequency	212,793

# Frequency per sample

	Frequency
Minimum frequency	5,583.0
1st quartile	7,626.0
Median frequency	8,997.0
3rd quartile	10,137.5
Maximum frequency	14,017.0
Mean frequency	9,251.869565217392

Frequency per sample detail (csv | html)

# Section 3: Build a phylogenetic tree

The next step does the following:

- 1. Perform an alignment on the representative sequences.
- 2. Mask sites in the alignment that are not phylogenetically informative.
- 3. Generate a phylogenetic tree.
- 4. Apply mid-point rooting to the tree.

A phylogenetic tree is necessary for any analyses that incorporates information on the relative relatedness of community members, by incorporating phylogenetic distances between observed organisms in the computation. This would include any beta-diversity analyses and visualisations from a weighted or unweighted Unifrac distance matrix.

```
mkdir analysis/tree
```

Use one thread only (which is the default action) so that identical results can be produced if rerun.

```
qiime phylogeny align-to-tree-mafft-fasttree \
--i-sequences analysis/dada2out/representative_sequences.qza \
--o-alignment analysis/tree/aligned_16s_representative_seqs.qza \
--o-masked-alignment
analysis/tree/masked_aligned_16s_representative_seqs.qza \
--o-tree analysis/tree/16s_unrooted_tree.qza \
--o-rooted-tree analysis/tree/16s_rooted_tree.qza \
--p-n-threads 1 \
--verbose
```

# Section 4: Basic visualisations and statistics

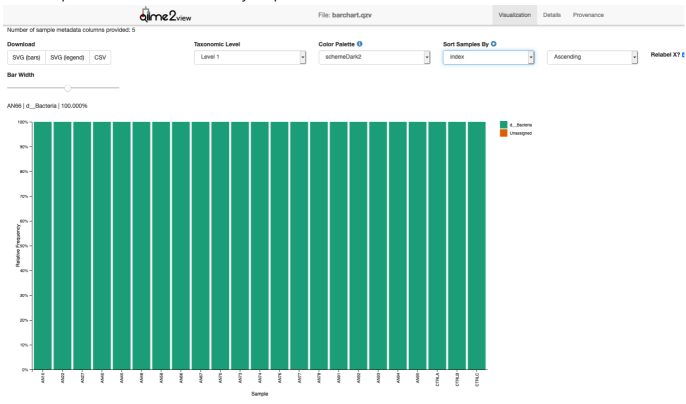
ASV relative abundance bar charts

Create bar charts to compare the relative abundance of ASVs across samples.

```
qiime taxa barplot \
 --i-table analysis/taxonomy/16s_table_filtered.qza \
 --i-taxonomy analysis/taxonomy/classification.qza \
 --m-metadata-file metadata.tsv \
 --o-visualization analysis/visualisations/barchart.qzv \
 --verbose
```

Copy analysis/visualisations/barchart.qzv to your local computer and view in QIIME 2 View (q2view). Try selecting different taxonomic levels and metadata-based sample sorting.

#### ??? example "Visualisations: Taxonomy Barplots"



```
![barplot2](./media/barplot_level3.png)
```

![barplot3](./media/barplot\_level5.png)

#### Rarefaction curves

Generate rarefaction curves to determine whether the samples have been sequenced deeply enough to capture all the community members. The max depth setting will depend on the number of sequences in your samples.

!!! info "Things to look for:" 1. Do the curves for each sample plateau? If they don't, the samples haven't been sequenced deeply enough to capture the full diversity of the bacterial communities, which is shown on the y-axis. 2. At what sequencing depth (x-axis) do your curves plateau? This value will be important for downstream analyses, particularly for alpha diversity analyses.

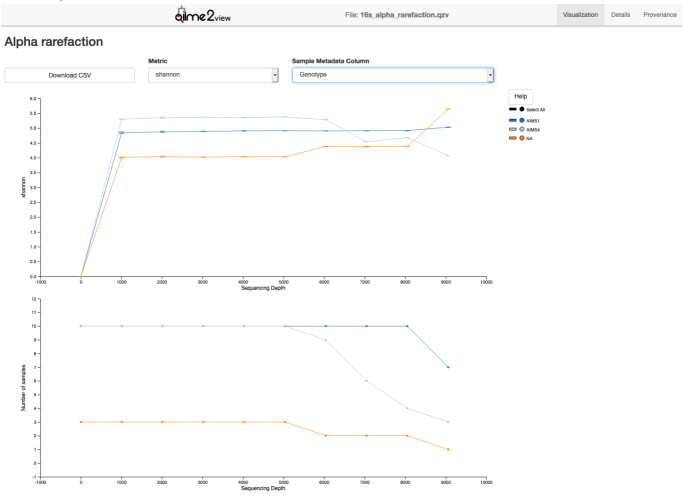
!!! note The value that you provide for --p-max-depth should be determined by reviewing the "Frequency per sample" information presented in the 16s\_table\_filtered.qzv file that was created above. In general, choosing a value that is somewhere around the median frequency seems to work well, but you may want to increase that value if the lines in the resulting rarefaction plot don't appear to be leveling out, or decrease that value if you seem to be losing many of your samples due to low total frequencies closer to the minimum sampling depth than the maximum sampling depth.

```
qiime diversity alpha-rarefaction \
--i-table analysis/taxonomy/16s_table_filtered.qza \
--i-phylogeny analysis/tree/16s_rooted_tree.qza \
```

```
--p-max-depth 9062 \
--m-metadata-file metadata.tsv \
--o-visualization analysis/visualisations/16s_alpha_rarefaction.qzv \
--verbose
```

Copy analysis/visualisations/16s\_alpha\_rarefaction.qzv to your local computer and view in QIIME 2 View (q2view).





## Alpha and beta diversity analysis

The following is taken directly from the Moving Pictures tutorial and adapted for this data set. QIIME 2's diversity analyses are available through the q2-diversity plugin, which supports computing alpha- and beta- diversity metrics, applying related statistical tests, and generating interactive visualisations. We'll first apply the core-metrics-phylogenetic method, which rarefies a FeatureTable[Frequency] to a user-specified depth, computes several alpha- and beta- diversity metrics, and generates principle coordinates analysis (PCoA) plots using Emperor for each of the beta diversity metrics.

The metrics computed by default are:

- Alpha diversity (operate on a single sample (i.e. within sample diversity)).
  - Shannon's diversity index (a quantitative measure of community richness)
  - Observed OTUs (a qualitative measure of community richness)

• Faith's Phylogenetic Diversity (a qualitative measure of community richness that incorporates phylogenetic relationships between the features)

- Evenness (or Pielou's Evenness; a measure of community evenness)
- Beta diversity (operate on a pair of samples (i.e. between sample diversity)).
  - Jaccard distance (a qualitative measure of community dissimilarity)
  - Bray-Curtis distance (a quantitative measure of community dissimilarity)
  - unweighted UniFrac distance (a qualitative measure of community dissimilarity that incorporates phylogenetic relationships between the features)
  - weighted UniFrac distance (a quantitative measure of community dissimilarity that incorporates phylogenetic relationships between the features)

An important parameter that needs to be provided to this script is —p—sampling—depth, which is the even sampling (i.e. rarefaction) depth that was determined above. As most diversity metrics are sensitive to different sampling depths across different samples, this script will randomly subsample the counts from each sample to the value provided for this parameter. For example, if —p—sampling—depth 500 is provided, this step will subsample the counts in each sample without replacement, so that each sample in the resulting table has a total count of 500. If the total count for any sample(s) are smaller than this value, those samples will be excluded from the diversity analysis. Choosing this value is tricky. We recommend making your choice by reviewing the information presented in the 16s\_table\_filtered.qzv file that was created above. Choose a value that is as high as possible (so more sequences per sample are retained), while excluding as few samples as possible.

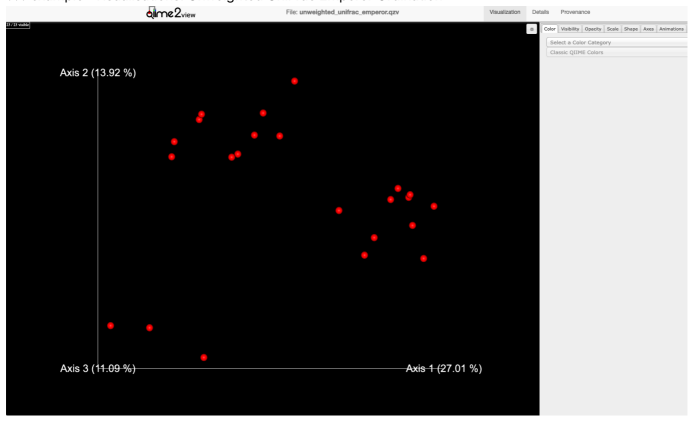
```
qiime diversity core-metrics-phylogenetic \
 --i-phylogeny analysis/tree/16s_rooted_tree.qza \
 --i-table analysis/taxonomy/16s_table_filtered.qza \
 --p-sampling-depth 5583 \
 --m-metadata-file metadata.tsv \
 --output-dir analysis/diversity_metrics
```

Copy the gaz files created from the above command into the visualisations subdirectory.

```
cp analysis/diversity_metrics/*.qzv analysis/visualisations
```

To view the differences between sample composition using unweighted UniFrac in ordination space, copy analysis/visualisations/unweighted\_unifrac\_emperor\_qzv to your local computer and view in QIIME 2 View (q2view).

#### ??? example "Visualisations: Unweighted UniFrac Emperor Ordination"



On q2view, select the "Colour" tab and the heading "Environment" in the dropdown menu and then by "Genotype" in the "Shape" tab.

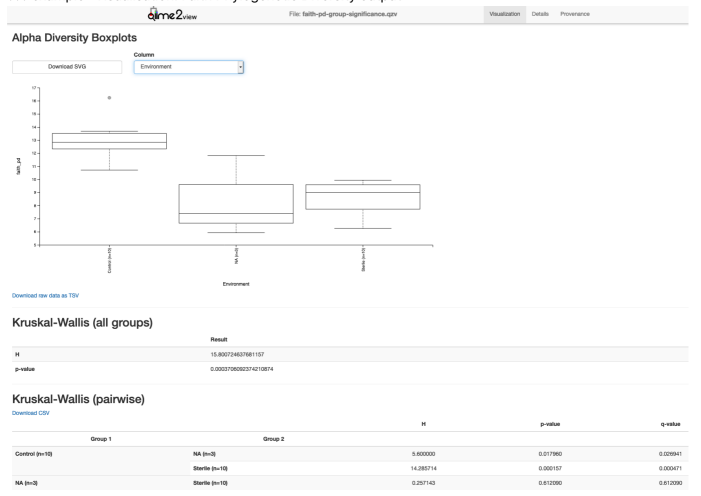
![unweighted\_unifrac\_emperor2](./media/unweighted\_unifrac\_emperor2.png)

Next, we'll test for associations between categorical metadata columns and alpha diversity data. We'll do that here for the Faith Phylogenetic Diversity (a measure of community richness) and evenness metrics.

```
qiime diversity alpha-group-significance \
 --i-alpha-diversity analysis/diversity_metrics/faith_pd_vector.qza \
 --m-metadata-file metadata.tsv \
 --o-visualization analysis/visualisations/faith-pd-group-
significance.qzv
```

Copy analysis/visualisations/faith-pd-group-significance.qzv to your local computer and view in QIIME 2 View (q2view).

## ??? example "Visualisation: Faith Phylogenetic Diversity output"



<sup>```</sup>python qiime diversity alpha-group-significance \ --i-alpha-diversity analysis/diversity\_metrics/evenness\_vector.qza \ --m-metadata-file metadata.tsv \ --o-visualization analysis/visualisations/evenness-group-significance.qzv ```

Copy analysis/visualisations/evenness-group-significance.qzv to your local computer and view in QIIME 2 View (q2view).

#### ??? example "Visualisation: Evenness output"

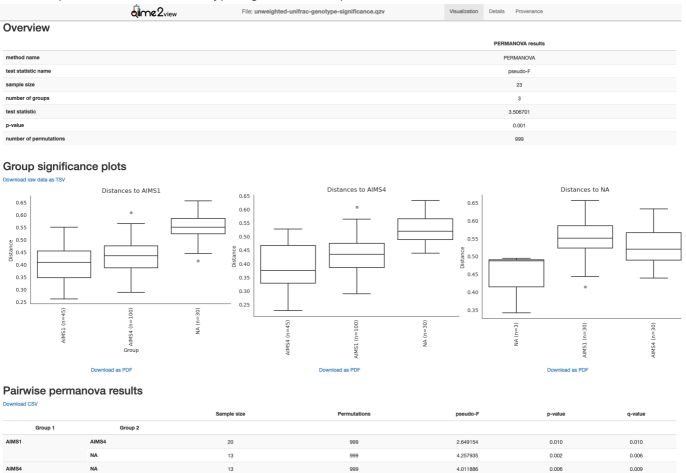


Finally, we'll analyse sample composition in the context of categorical metadata using a permutational multivariate analysis of variance (PERMANOVA, first described in Anderson (2001)) test using the betagroup-significance command. The following commands will test whether distances between samples within a group, such as samples from the same genotype, are more similar to each other then they are to samples from the other groups. If you call this command with the <code>--p-pairwise</code> parameter, as we'll do here, it will also perform pairwise tests that will allow you to determine which specific pairs of groups (e.g., AIMS1 and AIMS4) differ from one another, if any. This command can be slow to run, especially when passing <code>--p-pairwise</code>, since it is based on permutation tests. So, unlike the previous commands, we'll run beta-group-significance on specific columns of metadata that we're interested in exploring, rather than all metadata columns to which it is applicable. Here we'll apply this to our unweighted UniFrac distances, using two sample metadata columns, as follows.

```
qiime diversity beta-group-significance \
 --i-distance-matrix
analysis/diversity_metrics/unweighted_unifrac_distance_matrix.qza \
 --m-metadata-file metadata.tsv \
 --m-metadata-column Genotype \
 --o-visualization analysis/visualisations/unweighted-unifrac-genotype-significance.qzv \
 --p-pairwise
```

Copy analysis/visualisations/unweighted-unifrac-genotype-significance.qzv to your local computer and view in QIIME 2 View (q2view).

# ??? example "Visualisation: Genotype significance output"



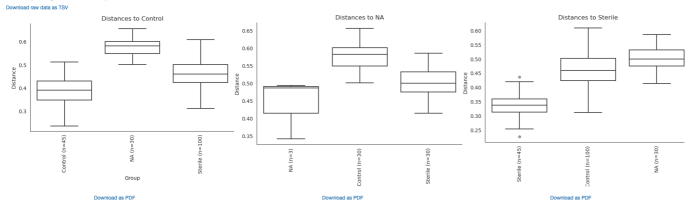
```python qiime diversity beta-group-significance \ --i-distance-matrix analysis/diversity\_metrics/unweighted\_unifrac\_distance\_matrix.qza \ --m-metadata-file metadata.tsv \ --m-metadata-column Environment \ --o-visualization analysis/visualisations/unweighted-unifrac-environment-significance.qzv \ --p-pairwise ```

Copy analysis/visualisations/unweighted-unifrac-environment-significance.qzv to your local computer and view in QIIME 2 View (q2view).

??? example "Visualisation: Environmental significance output"

	dime2view	File: unweighted-unifrac-environment-significance.qzv	Visualization Details Provenance
Overview			
			PERMANOVA results
method name			PERMANOVA
test statistic name			pseudo-F
sample size			23
number of groups			3
test statistic			5.896316
p-value			0.001
number of permutations			999

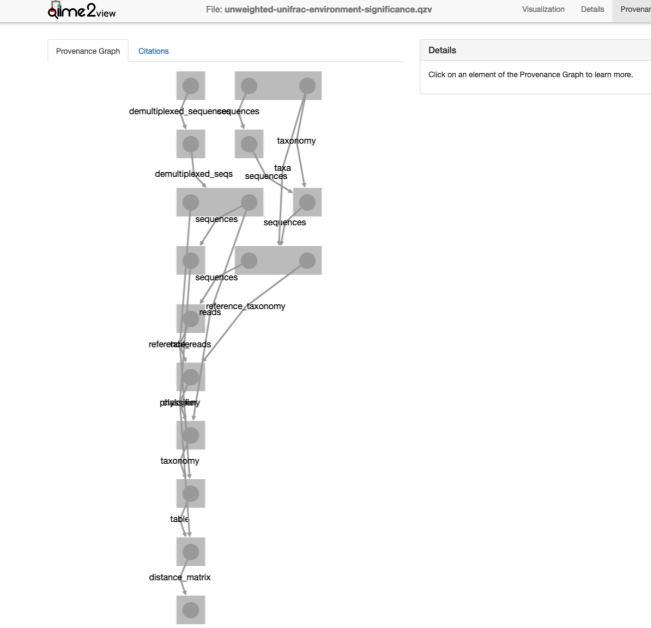
Group significance plots



Pairwise permanova results

Download COV		Sample size	Permutations	pseudo-F	p-value	q-value
Group 1	Group 2					
Control	NA	13	999	5.575155	0.002	0.003
	Sterile	20	999	6.895129	0.001	0.003
NA	Sterile	13	999	4.676336	0.009	0.009





Section 5: Exporting data for further analysis in R

You need to export your ASV table, taxonomy table, and tree file for analyses in R. Many file formats can be accepted.

Export unrooted tree as . nwk format as required for the R package phyloseq.

```
qiime tools export \
  --input-path analysis/tree/16s_unrooted_tree.qza \
  --output-path analysis/export
```

Create a BIOM table with taxonomy annotations. A FeatureTable[Frequency] artefact will be exported as a BIOM v2.1.0 formatted file.

```
qiime tools export \
    --input-path analysis/taxonomy/16s_table_filtered.qza \
    --output-path analysis/export
```

Then export BIOM to TSV

```
biom convert \
  -i analysis/export/feature-table.biom \
  -o analysis/export/feature-table.tsv \
  --to-tsv
```

Export Taxonomy as TSV

```
qiime tools export \
--input-path analysis/taxonomy/classification.qza \
--output-path analysis/export
```

Delete the header lines of the .tsv files

```
sed '1d' analysis/export/taxonomy.tsv >
analysis/export/taxonomy_noHeader.tsv
sed '1d' analysis/export/feature-table.tsv > analysis/export/feature-
table_noHeader.tsv
```

Some packages require your data to be in a consistent order, i.e. the order of your ASVs in the taxonomy table rows to be the same order of ASVs in the columns of your ASV table. It's recommended to clean up your taxonomy file. You can have blank spots where the level of classification was not completely resolved.